FOOD AND DRUG ADMINISTRATION (FDA)

Center for Drug Evaluation and Research (CDER)

Antimicrobial Drugs Advisory Committee (AMDAC) Meeting April 17, 2023

AGENDA

The committee will discuss new drug application (NDA) 216974, for sulbactam-durlobactam for injection, submitted by Entasis Therapeutics, Inc. The Applicant's proposed indication is treatment of hospital-acquired bacterial pneumonia (HABP) and ventilator-associated bacterial pneumonia (VABP) caused by susceptible strains of Acinetobacter baumannii-calcoaceticus complex (ABC) in adults.

9:00 a.m.	Call to Order	Lindsey R. Baden, MD Chairperson, AMDAC
9:10 a.m.	Introduction of Committee and Conflict of Interest Statement	Takyiah Stevenson, PharmD Acting Designated Federal Officer, AMDAC
9:15 a.m.	FDA Opening Remarks	Adam Sherwat, MD Deputy Director Office of Infectious Diseases (OID) Office of New Drugs (OND), CDER, FDA
9:25 a.m.	APPLICANT PRESENTATIONS	Entasis Therapeutics, Inc.
	Introduction	Shruta Rege, PhD Senior Vice President, Head of Regulatory Affairs and Development Operations Entasis Therapeutics
	Unmet Need	David Paterson, MBBS, PhD, FRACP Professor Saw Swee Hock School of Public Health National University of Singapore
	Microbiology and Pharmacology	Alita Miller, PhD Senior Vice President, Head of Research Entasis Therapeutics
	Efficacy	David Altarac, MD, MPA Chief Medical Officer Entasis Therapeutics

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AGENDA (cont.)

APPLICANT PRESENTATIONS

(CONT.)

Safety Drew Lewis, MD, MTM&H, FACP

Vice President, Clinical Development

Entasis Therapeutics

Clinical Perspective J. Patrik Hornak, MD

Assistant Professor of Medicine Division of Infectious Diseases

Assistant Clinical Director, AIDS Education &

Training Center

University of Texas Medical Branch at Galveston

Concluding Remarks Shruta Rege, PhD

Senior Vice President, Head of Regulatory Affairs

and Development Operations

Entasis Therapeutics

10:25 a.m. Clarifying Questions

10:45 a.m. **BREAK**

10:55 a.m. FDA PRESENTATIONS

Efficacy Assessment Karen Qi, PhD

Statistical Reviewer

Division of Biometrics IV

Office of Biostatistics, CDER, FDA

Clinical Safety Assessment Mayurika Ghosh, MD

Clinical Reviewer

Division of Anti-Infectives (DAI)

OID, OND, CDER, FDA

Clinical Microbiology Assessment Simone Shurland, PhD

Clinical Microbiology Reviewer DAI, OID, OND, CDER, FDA

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AGENDA (cont.)

Clinical Pharmacology	Xiaohui (Tracey) Wei, PhD
Assessment	Clinical Pharmacology Reviewer

Clinical Pharmacology Reviewer

Division of Infectious Disease Pharmacology Office of Clinical Pharmacology, CDER, FDA

11:55 a.m. Clarifying Questions

12:15 p.m. LUNCH

1:00 p.m. **OPEN PUBLIC HEARING**

2:00 p.m. Charge to the Committee Peter Kim, MD, MS

Director

DAI, OID, OND, CDER, FDA

Questions to the Committee/Committee Discussion 2:05 p.m.

2:52 p.m. **ADJOURNMENT**